

<https://helda.helsinki.fi>

Interaction of climate change with effects of conspecific and heterospecific density on reproduction

Moller, Anders Pape

2020-12

Moller , A P , Balbontin , J , Dhondt , A A , Adriaensen , F , Artemyev , A , Banbura , J , Barba , E , Biard , C , Blondel , J , Bouvier , J-C , Camprodon , J , Cecere , F , Charter , M , Cichon , M , Cusimano , C , Dubiec , A , Eens , M , Eeva , T , Ferns , P N , Forsman , J T , Goldshtein , A , Goodenough , A E , Gosler , A G , Gustafsson , L , Harnist , I , Hartley , I R , Heeb , P , Hinsley , S A , Jacob , S , Järvinen , A , Juskaitis , R , Korpimäki , E , Krams , I , Laaksonen , T , Leclercq , B , Lehtikoinen , E , Loukola , O , Mainwaring , M C , Mand , R , Massa , B , Matthysen , E , Mazgajski , T D , Merino , S , Mitrus , C , Monkkonen , M , Nager , R G , Nilsson , J , Nilsson , S G , Norte , A C , von Numers , M , Orell , M , Pimentel , C S , Pinxten , R , Priedniec , I , Remes , V , Richner , H , Robles , H , Rytönen , S , Senar , J C , Seppanen , J T , da Silva , L P , Slagsvold , T , Solonen , T , Sorace , A , Stenning , M J , Torok , J , Tryjanowski , P , van Noordwijk , A J , Walankiewicz , W & Lambrechts , M M 2020 , ' Interaction of climate change with effects of conspecific and heterospecific density on reproduction ' , Oikos , vol. 129 , no. 12 , pp. 1807-1819 . <https://doi.org/10.1111/oik.07305>

<http://hdl.handle.net/10138/333259>

<https://doi.org/10.1111/oik.07305>

unspecified

acceptedVersion

Downloaded from Helda, University of Helsinki institutional repository.

This is an electronic reprint of the original article.

This reprint may differ from the original in pagination and typographic detail.

Please cite the original version.

Interaction of climate change with effects of conspecific and heterospecific density on reproduction

Journal:	<i>Oikos</i>
Manuscript ID	OIK-07305.R2
Wiley - Manuscript type:	Research
Keywords:	Blue tit, great tit, interspecific competition, clutch size, laying date, temperature anomaly
Abstract:	<p>We studied the relationship between temperature and the coexistence of great tit <i>Parus major</i> and blue tit <i>Cyanistes caeruleus</i>, breeding in 75 study plots across Europe and North Africa. We expected an advance in laying date and a reduction in clutch size during warmer springs as a general response to climate warming and a delay in laying date and a reduction in clutch size during warmer winters due to density-dependent effects. As expected, as spring temperature increases laying date advances and as winter temperature increases clutch size is reduced in both species. Density of great tit affected the relationship between winter temperature and laying date in great and blue tit. Specifically, as density of great tit increased and temperature in winter increased both species started to reproduce later. Density of blue tit affected the relationship between spring temperature and blue and great tit laying date. Thus, both species start to reproduce earlier with increasing spring temperature as density of blue tit increases, which was not an expected outcome, since we expected that increasing spring temperature should advance laying date, while increasing density should delay it cancelling each other out. Climate warming and its interaction with density affects clutch size of great tits but not of blue tits. As predicted, great tit clutch size is reduced more with density of blue tits as temperature in winter increases. The relationship between spring temperature and density on clutch size of great tits depends on whether the increase is in density of great tit or blue tit. Therefore, an increase in temperature negatively affected the coexistence of blue and great tits differently in both species. Thus, blue tit clutch size was unaffected by the interaction effect of density with temperature, while great tit clutch size was affected in multiple ways by these interactions terms</p>

OIKOS

Research

- 0 **Interaction of climate change with effects of conspecific and** 61
heterospecific density on reproduction 65
- 5 Anders Pape Møller, Javier Balbontín, André A. Dhondt, Frank Adriaensen, Alexandr Artemyev, 70
 Jerzy Bańbura, Emilio Barba, Clotilde Biard, Jacques Blondel, Jean-Charles Bouvier, Jordi Camprodon,
 10 Francesco Cecere, Motti Charter, Mariusz Cichoń, Camillo Cusimano, Anna Dubiec, Marcel Eens,
 Tapio Eeva, Peter N. Ferns, Jukka T. Forsman, Aya Goldshtein, Anne E. Goodenough, Andrew G. Gosler,
 Lars Gustafsson, Iga Harnist, Ian R. Hartley, Philipp Heeb, Shelley A. Hinsley, Staffan Jacob,
 Antero Järvinen, Rimvydas Juškaitis, Erkki Korpimäki, Indrikis Krams, Toni Laaksonen, Bernard Leclercq,
 15 Esa Lehikoinen, Olli Loukola, Mark C. Mainwaring, Raivo Mänd, Bruno Massa, Erik Matthysen,
 Tomasz D. Mazgajski, Santiago Merino, Cezary Mitrus, Mikko Mönkkönen, Ruedi G. Nager,
 Jan-Åke Nilsson, Sven G. Nilsson, Ana C. Norte, Mikael von Numers, Markku Orell, Carla S. Pimentel,
 20 Rianne Pinxten, Ilze Priedniece, Vladimir Remeš, Heinz Richner, Hugo Robles, Seppo Rytkönen,
 Juan Carlos Senar, Janne T. Seppänen, Luís P. da Silva, Tore Slagsvold, Tapio Solonen, Alberto Sorace,
 Martyn J. Stenning, János Török, Piotr Tryjanowski, Arie J. van Noordwijk, Wiesław Walankiewicz and
 Marcel M. Lambrechts 85
- 25 *A. P. Møller* (<https://orcid.org/0000-0003-3739-4675>) ✉ (anders.moller@u-psud.fr), *Ecologie Systematique Evolution*, CNRS, AgroParisTech, Univ. Paris-Saclay, Orsay Cedex, France, and: Ministry of Education Key Laboratory for Biodiversity Science and Ecological Engineering, College of Life Sciences, Beijing Normal Univ., Beijing, PR China. – *J. Balbontín* (<https://orcid.org/0000-0003-1539-2636>), *Depto de Zoología, Facultad de Biología, Sevilla, Spain.* – *A. A. Dhondt*, *Laboratory of Ornithology, Cornell Univ., Ithaca, NY, USA.* – *A. V. Artemyev*, *Inst. of Biology of the Karelian Research Centre, Russian Academy of Sciences (IB KRC RAS), Russia.* – *J. Bańbura*, *Dept of Experimental Zoology and Evolutionary Biology, Univ. of Łódź, Łódź, Poland.* – *E. Barba*, *Terrestrial Vertebrates Research Unit 'Cavanilles', Inst. of Biodiversity and Evolutionary Biology, Univ. of Valencia, Paterna, Spain.* – *C. Biard*, *Sorbonne Univ., UPEC, Paris 7, CNRS, INRA, IRD, Inst. d'Écologie et des Sciences de l'Environnement de Paris, iEES Paris, Paris, France.* – *J. Blondel*, *Centre d'Ecologie Fonctionnelle et Evolutive, CEFE UMR 5175, Montpellier, France.* – *J.-C. Bouvier*, *INRA, UR 1115, Plantes et Systèmes de culture Horticoles, Avignon, France.* – *J. Camprodon*, *Àrea de Biodiversitat, Grup de Biologia de la Conservació, Centre Tecnològic Forestal de Catalunya, Solsona, Spain.* – *F. Cecere*, *Acquanegra sul Chiese (MM), Italy.* – *M. Charter*, *Shamir Research Inst., Univ. of Haifa, Katzrin, Israel.* – *M. Cichoń*, *Jagiellonian Univ., Inst. of Environmental Sciences, Krakow, Poland.* – *C. Cusimano*, *Dept of Agricultural and Forest Sciences, Univ. di Palermo, Italy.* – *A. Dubiec*, *I. Harnist and T. D. Mazgajski*, *Museum and Inst. of Zoology, Polish Academy of Sciences, Warsaw, Poland.* – *T. Eeva*, *E. Korpimäki*, *T. Laaksonen and E. Lehikoinen*, *Dept of Biology, Univ. of Turku, Turku, Finland.* – *P. N. Ferns*, *Cardiff Univ., School of Biosciences, Cardiff, S Glam, Wales.* – *J. T. Forsman*, *Natural Resources Inst. Finland, Oulu, Univ. of Oulu, Finland.* – *A. Goldshtein*, *Univ. of Haifa, Israel.* – *A. E. Goodenough*, *Dept of Natural and Social Sciences, Univ. of Gloucestershire, Cheltenham, UK.* – *A. G. Gosler*, *Dept of Zoology, Edward Grey Inst. of Field Ornithology and Inst. of Human Sciences, Oxford, UK.* – *L. Gustafsson*, *Dept of Animal Ecology, Evolutionary Biology Centre, Uppsala Univ., Uppsala, Sweden.* – *I. R. Hartley*, *Lancaster Environment Centre, Lancaster Univ., Lancaster, UK.* – *P. Heeb*, *Laboratoire Évolution & Diversité Biologique, UPS Toulouse III, Toulouse, France.* – *S. H. Hinsley*, *CEH Wallingford, Maclean Building, Crowmarsh Gifford, Wallingford, UK.* – *S. Jacob*, *Station d'Ecologie Théorique et Expérimentale, Moulis, France.* – *A. Järvinen*, *Kilpisjärvi Biological Station, Univ. of Helsinki, Finland.* – *R. Juškaitis*, *Inst. of Ecology of Nature Research Centre, Vilnius, Lithuania.* – *I. Krams and R. Mänd*, *Univ. Tartu, Inst. of Ecology and Earth Sciences, Tartu, Estonia.* – *B. Leclercq*, *Fleurey Sur Ouche, France.* – *O. Loukola*, *M. Orell and S. Rytkönen*, *Dept of Ecology and Genetics, Univ. of Oulu, Finland.* – *M. C. Mainwaring*, *Division of Biological Sciences, Univ. of Montana, Missoula, MT, USA.* – *B. Massa*, *Stazione Ornitologica, Palermo, Italy.* – *E. Matthysen*, *F. Adriaensen and M. Eens*, *Dept of Biology, Univ. of Antwerp, Antwerp, Belgium.* 90
 30 95
 35 100
 40 105
 45 110
 50 115
 55 121



- 0 – S. M. Rodríguez, CSIC, Depto de Ecología Evolutiva Museo Nacional de Ciencias Naturales, Agencia Estatal Consejo Superior de Investigaciones 61
Científicas CSIC, Madrid, Spain. – C. Mitrus, Dept of Vertebrate Ecology and Palaeontology, Wrocław Univ. of Environmental and Life Sciences,
Wrocław, Poland. – M. Mönkkönen, Univ. of Jyväskylä, Dept of Biological and Environmental Sciences, Univ. of Jyväskylä, Finland. – R. G. Nager,
Inst. of Biodiversity, Animal Health and Comparative Medicine, Univ. of Glasgow, Glasgow, UK. – J.-Å. Nilsson and S. G. Nilsson, Dept of Biology, Lund 65
Univ., Lund, Sweden. – A. C. Norte, MARE – Marine and Environmental Sciences Centre, Dept of Life Sciences, Largo Marquês de Pombal, Faculty of
Sciences and Technology, Univ. of Coimbra, Coimbra, Portugal. – M. von Numers, Environmental and Marine Biology, Åbo Akademi Univ. Åbo,
Finland. – C. S. Pimentel, Centro de Estudos Florestais, Instituto Superior de Agronomia, Univ. of Lisbon, Lisbon, Portugal. – R. Pinxten, Behavioural 70
Ecology and Ecophysiology Research Group, Dept of Biology, Wilrijk and Faculty of Social Sciences, Research Unit Didactica, Univ. of Antwerp, Antwerp,
Belgium. – I. Priedniece, Latvian Fund for Nature, Riga, Latvia. – V. Remeš, Laboratory of Ornithology, Dept of Zoology, Palacky Univ., Olomouc, Czech
Republic. – H. Richner, Univ. of Bern, Inst. of Ecology and Evolution (IEE), Bern, Switzerland. – H. Robles, Evolutionary Ecology Group (GIBE),
Faculty of Sciences, Univ. of A Coruña, A Coruña, Spain, and: Evolutionary Ecology Group (EVECO), Dept of Biology, Univ. of Antwerp, Antwerp, 75
Belgium. – J. C. Senar, Unidad de Ecología Evolutiva y de la Conducta, Museu de Ciències Naturals de Barcelona, Barcelona, Spain. – J.-T. Seppänen,
Univ. of Jyväskylä, Dept of Biological and Environmental Sciences, Univ. of Jyväskylä, Finland. – L. P. da Silva, CIBIO-InBIO, Research Center in
Biodiversity and Genetic Resources, Univ. of Porto, Vairão, Portugal. – T. Slagsvold, Dept of Biosciences, Univ. of Oslo, Oslo, Norway. – T. Solonen,
Luontotutkimus Solonen Oy, Helsinki, Finland. – A. Sorace, SROPU, Rome, Italy. – M. J. Stenning, School of Life Sciences, Univ. of Sussex, Sussex,
UK. – J. Török, Dept of Systematic Zoology and Ecology, Eötvös Loránd Univ., Budapest, Hungary. – P. Tryjanowski, Inst. of Zoology, Poznań Univ. of 80
Life Sciences, Poznań, Poland. – A. J. van Noordwijk, Netherlands Inst. of Ecology (NIOO-KNAW), Wageningen, the Netherlands. – W. Walankiewicz,
Siedlce Univ. of Natural Sciences and Humanities, Faculty of Natural Science, Dept of Zoology, Siedlce, Poland. – M. M. Lambrechts, Centre d'Ecologie
Fonctionnelle et Evolutive, CEFE UMR 5175, Campus CNRS, Montpellier, France.

20 We studied the relationship between temperature and the coexistence of great tit *Parus major* and blue tit *Cyanistes caeruleus*, 80
breeding in 75 study plots across Europe and North Africa. We expected an advance in laying date and a reduction in clutch
size during warmer springs as a general response to climate warming and a delay in laying date and a reduction in clutch size
during warmer winters due to density-dependent effects. As expected, as spring temperature increases laying date advances
and as winter temperature increases clutch size is reduced in both species. Density of great tit affected the relationship between 85
winter temperature and laying date in great and blue tit. Specifically, as density of great tit increased and temperature in winter
increased both species started to reproduce later. Density of blue tit affected the relationship between spring temperature and
blue and great tit laying date. Thus, both species start to reproduce earlier with increasing spring temperature as density of blue
tit increases, which was not an expected outcome, since we expected that increasing spring temperature should advance lay-
ing date, while increasing density should delay it cancelling each other out. Climate warming and its interaction with density 90
affects clutch size of great tits but not of blue tits. As predicted, great tit clutch size is reduced more with density of blue tits as
temperature in winter increases. The relationship between spring temperature and density on clutch size of great tits depends
on whether the increase is in density of great tit or blue tit. Therefore, an increase in temperature negatively affected the coex-
istence of blue and great tits differently in both species. Thus, blue tit clutch size was unaffected by the interaction effect of 95
density with temperature, while great tit clutch size was affected in multiple ways by these interactions terms.

Keywords: blue tit, clutch size, *Cyanistes caeruleus*, great tit, interspecific competition, intraspecific competition, laying date,
Parus major, temperature anomaly

Introduction

45 Climate change has been predicted to affect both intra- and interspecific competition either through effects on the abun-
dance of limiting resources, through changes in the abun-
dance of interacting species or through changes in species
distribution (Møller et al. 2010, 2019). It is well known that
density-dependent effects on fecundity or other demographic
50 traits can regulate populations as a result of intra- and/or
interspecific competition. There are two mechanisms that
explain density-dependence in fecundity. First, as density
increases breeding habitats are occupied in sequential order
of quality. High quality sites are occupied firsts and poor
55 quality sites later (i.e. those that yield lower than average
fecundity) resulting in a decrease of reproductive parameters
at the population level (i.e. Habitat heterogeneity hypoth-
esis) (Dhondt et al. 1992). This hypothesis can be extended
60 to the individual level whereby high quality or older indi-
viduals settle first and hence occupy the high quality sites
while low quality individuals appear later and occupy low
quality sites. This also would enhance the decrease in repro-
ductive parameters at the population level (Balbontín and
105 Ferrer 2008). Second, increased density can also reduce
reproductive parameters due to antagonistic encounters
amongst individuals (i.e. Interference hypothesis) (Dhondt
and Schillemans 1983) or through competition for food. In
birds, advanced phenology is correlated with increased fecun-
110 dity (Kluijver 1951, Winkler and Allen 1996, Smith and
Moore 2004). Thus, it is expected that competition caused
by increased density would result in delayed breeding at the
population level and hence reduce fecundity (e.g. clutch size).
Intra- and interspecific competition are known to affect lay-
115 ing date and clutch size in hole-nesting birds (Dhondt 2010,
2012, Stenseth et al. 2015, Møller et al. 2018). However,
some studies suggest that such effects of competition are
significant only in specific plots or specific periods, raising
120 questions about the generality of these phenomena, but also

0 about their underlying causes (Alatalo and Lundberg 1984,
Török and Tóth 1988, Dhondt et al. 1992).

5 Birds advance the timing of migration or breeding in
response to climate warming (Parmesan and Yohe 2003).
This response has been attributed to improve the synchro-
nization of the timing of reproduction with the timing of
prey emergence at lower trophic levels. Thus, the peak in
food abundance (e.g. caterpillar) would match with the peak
of maximum demand of chicks during the nestling stage.
10 However, this possible mismatch could vary across Europe
(Visser et al. 1998, 2009). Although the relationship between
laying date and climate warming is well known, we know
little about the relationship between clutch size and temper-
ature. For instance, a warmer spring could provide a more
favourable ambient environment for females that might
result in an increase in reproductive investment. However, it
has been suggested that a reduction in clutch size could be an
alternative strategy to improve reproductive adjustment with
lower trophic levels (Bleu et al. 2017).

20 Female great and blue tits lay a clutch of 7–9 eggs and
7–13 eggs, respectively (Perrins 1991). The species differ in
life-history strategies in that the probability to lay a second
brood is lower in blue compared to great tit (Gibb and Betts
1963, Visser et al. 2003). As double-brooded species have
more difficulty to cope with climate change than single-
brood species (Husby et al. 2009) we expected a difference in
response to climate change of blue tits compared to great tits.
Blue tit should respond more strongly to increasing tempera-
tures since it is a mainly single-brood species.

30 During the breeding season interspecific competition for
food occurs when the smaller blue tit consumes smaller instar
of the same caterpillar species as eaten by the larger great tit.
This results in pre-emptive food consumption that differ-
entially impairs the great tit, which is therefore the inferior
competitor (Dhondt 2012). On the other hand, competition
for access to nest boxes favours great tits when large-holed
nest boxes are available (Dhondt and Eyckerman 1980).
40 Great tits outcompete blue tits by excluding blue tits from
boxes used as roosting or breeding sites, even when boxes are
super-abundant (Dhondt 2012).

During the non-breeding season great and blue tit adult
survival rate is affected by winter severity (Robinson et al.
45 2007, Pearce-Higgins and Green 2014). Winters with abun-
dant and extended snow cover decrease survival rate of adult
and one-year old birds and consequently population size is
reduced during the next breeding season. These effects of
winter weather on survival have been attributed to reduced
food supply in snowy winters (e.g. beech mast) (Pearce-
Higgins and Green 2014 and reference herein). Thus, we
expected an increase in population size (i.e. density) after
warmer winters. Specifically, as density increased, the propor-
tion of one-year old individuals being part of the popula-
55 tions this effect should specifically affect short-lived species.
This could constitute part of the mechanism that could delay
laying date and reduce clutch size due to density-dependent
effects (Dhondt et al. 1992, Balbontín and Ferrer 2008).

60

Recently, Møller et al. (2018) published extensive analyses
of effects of competition on laying date and clutch size in
great and blue tit across Europe and North Africa. Here we
expand these analyses by testing for the first time whether the
effects of climate change during the last 50 years significantly
influenced intra- and interspecific competition by measur-
ing their presumed effect on lay date and clutch size while
controlling statistically for a number of variables that are
known to predict lay date and clutch size in blue and great
tit (Møller et al. 2014a). We did so by analysing a long-term
dataset of 75 studies of two species of secondary hole nesting
birds across Europe and North Africa. We used density (num-
ber of occupied nestboxes per ha) as a proxy for intensity of
competition (Dhondt 2012). Intensity of competition could
be measured as the slope of density on a demographic param-
eter (e.g. laying date or clutch size). Therefore, we included
density as an independent variable in statistical models where
the dependent variable was laying date or clutch size to mea-
sure intensity of competition (Welden and Slauson 1986).
We used temperature anomalies during winter and spring as
a proxy of climate change.

The objectives of this study are to test whether laying date
advanced and clutch size change with increasing temperature,
whether this effect was modified by density; and whether the
intensity of intra- and interspecific competition was impacted
by increasing temperature anomalies.

We predicted that 1) in warmer winters survival rates
should increase, particularly for yearling birds (i.e. those that
lay smaller clutches and lay later) in both great and blue tits.
Hence inter- and intraspecific density should increase result-
ing in a delayed laying date and a reduced clutch size. The
relationships between temperature anomalies in winter and
laying date and clutch size are predicted to increase with den-
sity. Thus, we expected a greater delay in laying date and a
greater reduction in clutch size with increased winter temper-
ature anomalies (for details see Table 1). 2) In warmer springs
(i.e. with increased spring temperature anomalies) a more
proximal cue for seasonally breeding species, we expected lay-
ing date to advance in both great and blue tit, with a stron-
ger response in blue tits. If this results in a mismatch with
lower trophic levels, tits should adaptively reduce clutch size
to cope with the reduced food availability when feeding nest-
lings. We thus expect that positive spring temperature anom-
alies will lead to a reduced clutch size with a stronger effect
in great tits. We did not predict an effect for the outcome of
the interaction between temperature anomalies in spring and
density on laying date. Because warming should advance lay-
ing, and competition should delay laying, both effects could
cancel each other out. Likewise, competition and its inter-
action with temperature anomalies in spring should reduce
clutch size in great tits more than blue tits. This is expected
because the latter outcompetes the former for food, and also
because blue tits should respond more rapidly to tempera-
tures in spring since it is mainly a single-brood species (for
details see Table 1).

110
115
120

Table 1. Predictions from hypotheses about response to climate warming and its interaction with density in great and blue tits studied across 75 study sites in Europe and North Africa. The predictions are depicted as expected from statistical models described in Methods and presented in Table 2–5. GT is great tit. BT is blue tit. T^a = temperature anomaly. Nfloor = nest floor surface (mm). Models were run separately for GT and BT laying date and clutch size, respectively. Full models are in Table foot. Symbol| refers to the random effects part of the formula. Pred. is prediction number. Model formula only depicted the variables of interest related to the predicted relationship.

Pred.	Model formula	Predict relationship
1	Laying date (GT) $\sim \beta_0 + \beta_1 \times T^a$ spring Laying date (BT) $\sim \beta_0 + \beta_2 \times T^a$ spring	$\beta_1 \neq 0$ and $\beta_1 < 0$. Laying date advance as T ^a spring increases $\beta_2 \neq 0$ and $\beta_2 < 0$. Idem. $\beta_2 > \beta_1$. BT advance more laying date than GT as T ^a spring increases
2	Laying date $\sim \beta_0 + \beta_1 \times T^a$ winter	$\beta_1 \neq 0$ and $\beta_1 > 0$. Laying date delayed as T ^a winter increases
3	Laying date $\sim \beta_0 + \beta_7 \times (T^a \text{ winter: Density})$	$\beta_7 \neq 0$ and $\beta_7 > 0$. Laying date delayed more with density as T ^a winter increases
4	Laying date $\sim \beta_0 + \beta_7 \times (T^a \text{ spring: Density})$	$\beta_7 = 0$ No effect
5	Clutch size (GT) $\sim \beta_0 + \beta_1 \times T^a$ spring Clutch size (BT) $\sim \beta_0 + \beta_2 \times T^a$ spring	$\beta_1 \neq 0$ and $\beta_1 < 0$. Clutch size is reduced as T ^a spring increases $\beta_2 \neq 0$ and $\beta_2 < 0$. Idem. $\beta_1 < \beta_2$. GT reduced more clutch size than BT as T ^a spring increases
6	Clutch size $\sim \beta_0 + \beta_1 \times T^a$ winter	$\beta_1 \neq 0$ and $\beta_1 < 0$. Clutch size is reduced as T ^a spring increases
7	Clutch size $\sim \beta_0 + \beta_7 \times (T^a \text{ winter: Density})$	$\beta_7 \neq 0$ and $\beta_7 < 0$. Clutch size is reduced more with density as T ^a winter increases
8	Clutch size $\sim \beta_0 + \beta_7 \times (T^a \text{ spring: Density})$	$\beta_7 \neq 0$ and $\beta_7 < 0$. Clutch size is reduced more with density as T ^a spring increases
Laying date $\sim \beta_0 + \beta_1 \times T^a \text{ spring} + \beta_2 \times T^a \text{ winter} + \beta_3 \times \text{Density GT} + \beta_4 \times \text{Density BT} + \beta_5 \times (T^a \text{ spring: Density GT}) + \beta_6 \times (T^a \text{ spring: Density BT}) + \beta_7 \times (T^a \text{ winter: Density GT}) + \beta_8 \times (T^a \text{ winter: Density BT}) + \beta_9 \times \text{Latitude} + \beta_{10} \times \text{Longitude} + \beta_{11} \times \text{Altitude} + \beta_{12} \times \text{Habitat} + \beta_{13} \times \text{Urbanization} + \beta_{14} \times \text{Nfloor} + \beta_{15} \times \text{Nest material} + (1 \text{Site}) + (1 \text{Year})$.		
Clutch size $\sim \beta_0 + \beta_1 \times T^a \text{ spring} + \beta_2 \times T^a \text{ winter} + \beta_3 \times \text{Density GT} + \beta_4 \times \text{Density BT} + \beta_5 \times (T^a \text{ spring: Density GT}) + \beta_6 \times (T^a \text{ spring: Density BT}) + \beta_7 \times (T^a \text{ winter: Density GT}) + \beta_8 \times (T^a \text{ winter: Density BT}) + \beta_9 \times \text{Laying date} + \beta_{10} \times \text{Latitude} + \beta_{11} \times \text{Longitude} + \beta_{12} \times \text{Altitude} + \beta_{13} \times \text{Habitat} + \beta_{14} \times \text{Urbanization} + \beta_{15} \times \text{Nfloor} + \beta_{16} \times \text{Nest material} + (1 \text{Site}) + (1 \text{Year})$.		

Methods

Study sites and data sets

We obtained information on density of occupied nest boxes per ha, nest box size, clutch size, laying date and ecological variables from all studies considered in this paper for two common species of secondary hole-nesters, the great tit and the blue tit, across Europe and North Africa, as described in detail elsewhere (Møller et al. 2014a, b). Density of great or blue tits was estimated as the number of occupied nest boxes per ha. Nest boxes have entrance holes that could be large enough for great tits (32 mm diameter) and in some plots have additional small-holed nest boxes available for blue tits (26 mm diameter). Nest boxes usually are available in all plots at high densities (e.g. $> 6.6 \text{ ha}^{-1}$) and did not limit population size. In total, we calculated 919 yearly mean laying dates and 916 yearly mean clutch sizes across 75 study plots with both great and blue tits breeding during the period 1957–2012 (Møller et al. 2014a, b). Study years started in 1957 in Vlieland and ended in 2012 in several study plots. The mean (SD) numbers of years monitored was 11.49 (14.66) (range: 1–55 years). The mean (SD) nearest neighbour distance was 173.6 (115.4) km.

The abundance of great and blue tits changes across years within study plots. The average abundance of great tits was 18 (range: 1–137) occupied nest boxes and for blue tits 17 (range: 1–99) occupied nest boxes. The average density was 0.68 breeding pairs/ha for great tit and 0.65 breeding pairs/ha for blue tit. We included only study plots in which both great and blue tits had been recorded breeding at least once

in order to ensure that all study plots contained suitable habitats, breeding sites and nest boxes for both species. Although the taxonomy of tits is currently under revision (Stenning 2018), we used two taxa of tits (great tit and blue tit) with comparable ecologies without considering that some populations in the Canary Islands and North Africa may constitute the separate species *Cyanistes teneriffae* (Stenning 2018).

We restricted the analyses to first clutches, or early clutches known to be initiated less than 30 days after the first egg was laid in a given year in a study area to standardize sampling procedures (cf. Nager and van Noordwijk 1995). We assumed that the very small number of unidentified early repeat clutches that usually result from perturbations (Haywood 1993), or lay-dates calculated from information obtained from different breeding stages (Lambrechts et al. 1997), did not substantially alter the overall average clutch size and average lay date per study plot. Second or late clutches were excluded from analyses because they are usually smaller than first or early clutches and their frequency varies between years and habitats (Kluijver 1951, Lambrechts et al. 2008).

All data are available at: doi:10.5061/dryad.p763611.

Life-history traits and environmental factors

Information on latitude, longitude and altitude was provided by the authors of earlier studies or found in publications (for details see Møller et al. 2014a, b, 2018, Vaugoyeau et al. 2016). Tree species vary significantly in timing and amount of invertebrates available for raising offspring in tits (Kennedy and Southwood 1984, Lambrechts et al. 2008). We classified vegetation as ‘deciduous’ habitat dominated by non-evergreen

0 broad-leaved deciduous trees (*Alnus*, *Betula*, *Carpinus*, *Citrus*,
1 *Fagus*, *Fraxinus*, *Malus*, *Quercus*, including *Q. faginea*), ‘ever-
2 green’ habitat dominated by non-coniferous broad-leaved
3 evergreen trees (*Q. ilex*, *Q. suber*), ‘coniferous’ habitat domi-
4 nated by coniferous trees (*Abies*, *Cedrus*, *Picea*, *Pinus*), or
5 ‘mixed’ habitats formed by a combination of the former tree
6 vegetation classes (e.g. deciduous mixed with evergreen).
7 Study plots were classified as either rural or urban, using the
8 classifications provided by the original studies. Urban areas
9 were characterized by city parks, gardens and similar habitats
10 in close proximity to humans, while forests, plantations and
11 similar habitats were classified as rural. Only nest box studies
12 were included. We calculated the internal floor area (in cm²)
13 of nest boxes, using publications (Lambrechts et al. 2010) or
14 additional information provided by participants. The materi-
15 al constituting the nest box was divided into two broad
16 binary classes that are readily distinguishable: wood scored
17 as 1, which includes tree trunks, plywood, board-masonite or
18 board (Gustafsson and Nilsson 1985) and concrete scored as
19 0 (a mixture of cement and other materials; Lambrechts et al.
20 2010). We included all these input variables in the statistical
21 models because previous studies have indicated that each of
22 these variables are significant predictors of laying date and
23 clutch size (Lambrechts et al. 2010, Møller et al. 2014a, b,
24 Vaugoyeau et al. 2016).

Major life history traits are known to vary among years. For
instance, in local study plots biotic (e.g. resource availability,
intra- or interspecific interactions) and abiotic factors (e.g.
weather) can vary substantially among years perhaps explain-
ing within-plot variation in average clutch size (Kluijver
1951, Perrins 1965, Both 2000). We thus used study year as
a random factor in all analyses.

35 Temperature

We used temperature anomalies rather than temperatures
themselves to characterize climate across the very large geo-
40 graphic area of the study because we were interested in the
effect of temperature change at each study site. Temperature
anomaly is defined as a departure from a reference value or
long-term average. A positive anomaly indicates that the
observed temperature was warmer than the reference value,
45 while a negative anomaly indicates that the observed temper-
ature was cooler than the reference value. If, for example, the
reference value is 15°C and the measured temperature is 17°C,
then the temperature anomaly is +2°C (17–15°C (CMB and
Crouch 2012). Mean great tit laying date was 22 April (range:
50 19 March–12 May, n=75 study plots) and mean blue tit
laying date was 23 April (range: 17 March–13 May, n=75
study plots). Thus, we selected mean spring temperature
anomalies during March–May as an appropriate time win-
55 dow that would reflected the temperatures experienced prior
to the start of reproduction for all plots. Likewise, we used
the mean winter temperature anomalies during December–
February and mean annual temperature anomalies estimated
as the temperature anomaly for every year. These tempera-
60 ture anomalies were calculated with respect to the average

temperature obtained for the time window 1980–2010 taken
for each study plot. Temperature anomalies for each study
plot were extracted from a 5 × 5 degree-grid where tempera-
61 ture anomalies were calculated by averaging the anomalies for
each meteorological station that is found within a grid point
65 (<www.ncdc.noaa.gov/cag/>). Temperature anomalies tend
to be highly positively correlated across distances of less than
1000’s km (Hansen and Lebedeff 1987) and hence using a 5
70 × 5 degree grid was adequate.

Statistical analyses

We tested whether temperature anomaly interacts with den-
sity affecting the competitive or coexistence relationship
between great and blue tits to affect laying date and clutch
size by evaluating possible candidate models obtained from
four general linear mixed models. We evaluated predictors
explaining four response variables (i.e. laying date of great
and blue tits and clutch size of great and blue tits, respec-
80 tively) by defining for each response a full model. The models
were set with the aim to test the predictions stated in Table 1
(see full model formula in footnotes). Each of these full mod-
els included the density of great and blue tits and tempera-
85 ture anomaly of spring and winter. We explicitly included
the interaction term between (winter or spring) temperature
anomaly and density of great or blue tits to test the hypoth-
esis that an increase in temperature anomaly could affect lay-
ing date or clutch size of either species. We also included the
90 previously listed confounding variables that have been shown
to affect laying date or clutch size of great and blue tits. These
input variables were latitude, longitude, altitude, nest box
material (wood or concrete), nest floor surface, urbanization
(rural or urban) and habitat (coniferous, deciduous, mixed or
95 evergreen). The full model was a linear mixed effect model in
which we included these predictors as fixed effects and two
random intercepts for study plot and year that were retained
in all models. We included in the same model density of blue
100 and great tit and temperature anomaly in spring and win-
ter because these variables were only moderately positively
correlated (density of great tit versus density of blue tit:
 $r_s=0.441$, $p < 0.001$; temperature anomaly in spring versus
temperature anomaly in winter: $r_s=0.497$, $p < 0.001$). We
105 calculated variance inflation factors (VIF) in the four models
(Freund et al. 2003). All predictors showed low VIF values
less than 5.0.

For all models we first standardized the input variables
entering the full model by scaling them after centering their
mean and dividing by two standard deviations, which allowed
comparison on the same scale of coefficients of binary factors
and covariates. Therefore, the parameter estimates were stan-
dardized effect sizes on a comparable scale (Gelman 2008,
Grueber et al. 2011). In total, for the analyses on laying date
we included in each full model 15 predictor terms result-
115 ing in $2^{15}=32\,768$ candidate models. For the analyses on
clutch size we also included laying date in each full model
which resulted in 16 terms or $2^{16}=65\,536$ candidate mod-
els. Top models of all possible candidates were determined
121

using Akaike information criterion with a correction for small sample size (AICc). We calculated Akaike weight (w) for each candidate model that can be interpreted as the probability that it is the best model, given the data and set of candidate models (Burnham and Anderson, 2002). The reference level of the fixed factor 'habitat' was 'conifer', and for 'urbanization' it was 'rural' and for 'material' it was 'concrete' in all models. The final model was obtained by averaging the parameter estimates from top models at a cut-off criterion of AIC < 6.0 (Richards 2008). We employed the package MuMIn (Bartón 2009) and the package lme4 (Bates and Maechler 2009) using R ver. 3.3.2 <www.r-project.org>. The confidence intervals (CI 95%) were calculated from the final model using the parameter estimates (effect size) and the associated SE obtained after model averaging. We assumed that a predictor term significantly contributed to explaining the response variable when the CI for the estimated parameter excluded zero (Grueber et al. 2011).

Results

Temporal trend in temperature anomaly

The annual temperature anomaly (January–December) increased on average $+0.027^{\circ}\text{C}/\text{year}$ ($F=307.4$, $\text{df}=1$, 462 , $p < 0.0001$, estimate (SE) = 0.027 (0.001)). Spring temperature anomaly for March–May increased on average by $+0.033^{\circ}\text{C}/\text{year}$ ($F=202.60$, $\text{df}=1$, 462 , $p < 0.0001$, estimate (SE) = 0.033 (0.002)). Winter temperature anomaly for December–February increased on average by $+0.011^{\circ}\text{C}/\text{year}$ ($F=12.06$, $\text{df}=1$, 462 , $p < 0.0006$, estimate (SE) = 0.011 (0.003)). These results were obtained for three different linear mixed-effect models where study plot was included as a random term with temperature anomaly (year, winter or spring) as the dependent variable and year (centered as its overall mean) included as a fixed effect. The inclusion of a random slope for year within study plot did not improve the models (e.g. analyses where models with and without a random slope for year: Temperature anomaly (year): Likelihood-ratio = 0.078 , $p=0.96$; Temperature anomaly (spring): L-ratio = 0.000 , $p=1.0$; Temperature anomaly (winter): L-ratio = 0.000 , $p=1.0$), and, therefore, a common slope for year was assumed in these analyses. Thus, the increase in temperature anomaly did not differ significantly amongst study plots.

Inter- and intraspecific competition and response to climate change

Great tit timing of reproduction

The models of the analyses for great tit showed that laying date advanced with increasing spring temperature anomaly (Table 2, Fig. 1). The interaction term between spring temperature anomaly and blue tit density, on the one hand, and winter temperature anomaly and great tit density, on the other, also influenced great tit laying date significantly.

Table 2. Summary results after model averaging of the effects of temperature anomaly in spring and winter and density of great and blue tits on laying date of great tits across Europe and North Africa. Estimates (SE) and 95% CI are reported. The model also included other potential predictors of laying date of great tit. Sample size is 919 observations on yearly average laying date taken across 75 sites across Europe. Effects in bold font are statistically significant.

Parameter	Estimate	SE	CI (95%)
Intercept	58.910	2.130	(54.729, 63.091)
Density of blue tit (BT)	1.497	0.766	(−0.006, 3.002)
Density of great tit (GT)	−1.730	0.611	(−2.928, −0.530)
Latitude	8.766	2.738	(3.392, 14.139)
Longitude	5.295	1.918	(1.531, 9.060)
Floor surface (nest)	0.069	1.145	(−2.177, 2.317)
T^a Spring	−5.100	0.626	(−6.330, −3.870)
T ^a winter	−0.977	0.559	(−2.075, 0.121)
T^a Spring × BT	−2.452	0.849	(−4.199, −0.875)
T ^a Spring × GT	−1.267	0.760	(−2.759, 0.224)
T ^a Winter × BT	−0.521	0.913	(−2.313, 1.270)
T^a Winter × GT	1.535	0.702	(0.157, 2.914)
Altitude	0.389	2.577	(−4.668, 5.448)
Material	2.027	1.837	(−1.577, 5.632)
Urbanization	−2.553	1.259	(−5.026, −0.081)
Habitat (Deciduous)	−4.858	2.461	(−9.689, −0.027)
Habitat (Evergreen)	1.395	3.603	(−5.677, 8.467)
Habitat (Mixed)	−3.432	1.039	(−5.471, −1.392)

* Effect sizes have been standardized to two SD following Gelman (2008). Reference levels were (no) urbanized for urbanization and (conifer) for habitat. Parameter estimates for predictors with confidence intervals not including zero are highlighted in bold. CI is confidence interval.

Thus, great tits laid earlier as spring temperature anomaly increased and laying date advanced more steeply with spring temperature anomaly as blue tit density increased (Fig. 3a). In contrast, great tit laying date was delayed with winter temperature anomaly as great tit density increased (Fig. 3b). These effects were found while controlling for confounding variables known to affect laying date, such as latitude, longitude, altitude, habitat, urbanization, nest floor surface and nest-box material.

Blue tit timing of reproduction

The final model for blue tit laying date was similar to that for great tits. Across Europe and North Africa blue tits advanced laying date as spring temperature anomaly increased (Table 3, Fig. 1). Similarly to the results in great tits, blue tit lay date was significantly correlated with the interaction between spring temperature anomalies and heterospecific density, and in this case it was also correlated with the interaction between spring temperature anomalies and conspecific density. Likewise, blue tit laying date was significantly correlated

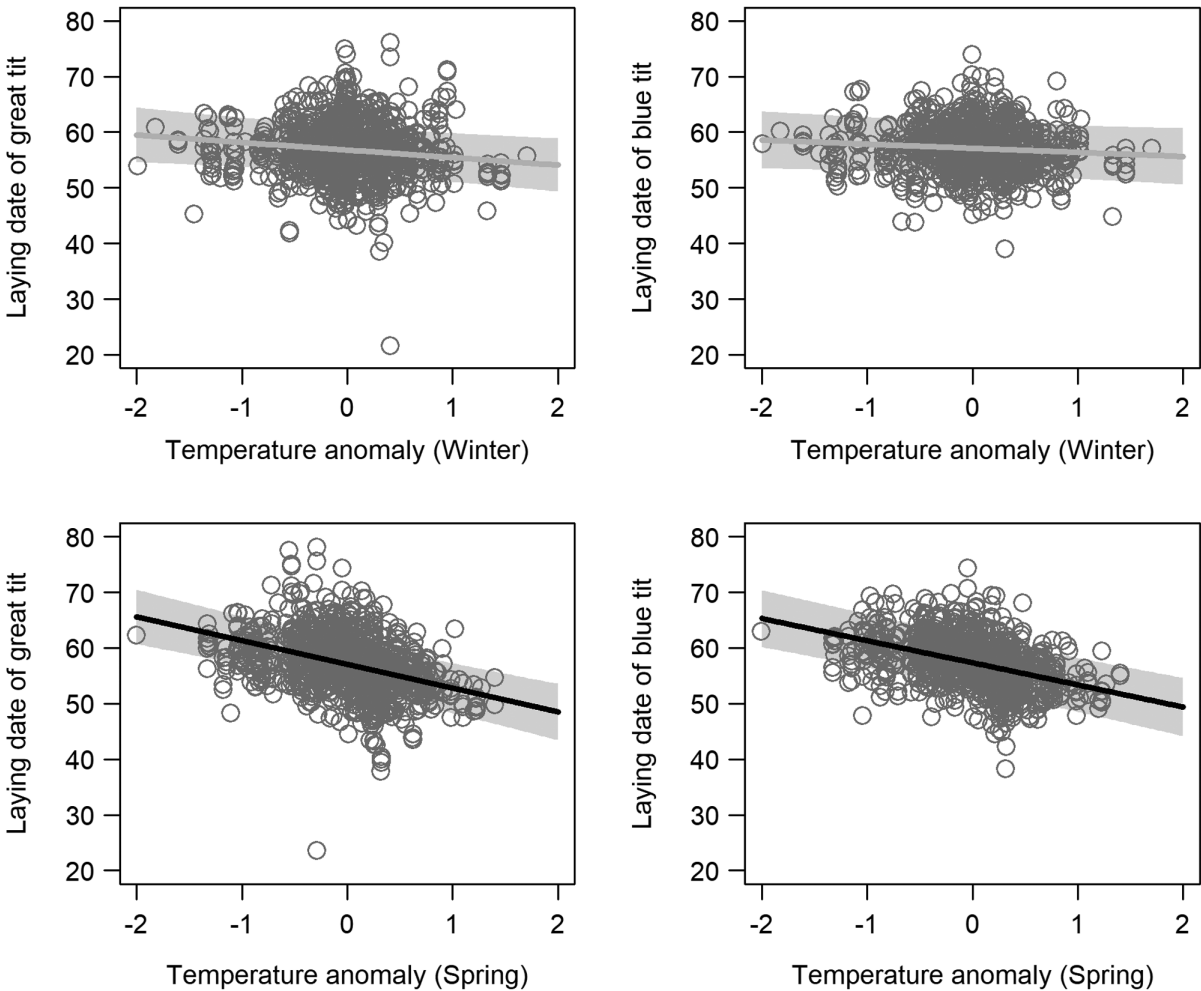


Figure 1. Effects of temperature anomaly (°C) in spring and winter on laying date of great and blue tits where 1 = 1 January. The lines (\pm 95% CI) are the predicted values obtained from a linear mixed-effect model where latitude, longitude and altitude and nest floor surface are maintained at their mean values and habitat, urbanization and nest box material at their reference values (i.e. conifer, rural habitat and box material concrete). Black lines represent effect size and did not include zero in 95% CI and grey lines represent effect size and did include zero in 95% CI (see Table 3 and 4 for details).

with the interaction of great tit density with winter temperature anomaly. Thus, blue tits laid earlier as spring temperature anomalies increased and laying date advanced more steeply with spring temperature anomaly as blue tit density increased, this effect being modulated by the interaction of temperature anomaly both with great and blue tit density (e.g. Fig. 4a for the interaction between density of blue tit and spring temperature anomaly on laying date of blue tit). In contrast, blue tit laying date was delayed more steeply with winter temperature anomaly as density of great tits increased (Fig. 4b). This was the case for the model analysing great tit laying date (Fig. 3b). These effects were found while controlling for confounding variables known to affect laying date, such as latitude, longitude, altitude, habitat, urbanization, nest floor surface and nest-box material.

Effects on great tit clutch size

Across Europe and North Africa great tit clutch size decreased as winter temperature anomaly increased, and it was not correlated with the main effect of spring temperature anomaly (Table 4, Fig. 2). We found opposite interaction effects between temperature anomaly in spring and winter and the density of blue tits on clutch size of great tits. We also found opposite effects (i.e. different sign) on the interaction between spring temperature anomaly and density of great or blue tit on great tit clutch size. Thus, there was a negative interaction between spring temperature anomaly and great tit density on great tit clutch size, and a positive interaction between effect of spring temperature anomaly and blue tit density on great tit clutch size. Thus, great tit clutch size was further reduced with increasing spring temperature anomaly as great tit density increased, and with increasing winter temperature anomaly as blue tit density increased (Fig. 5a–b).

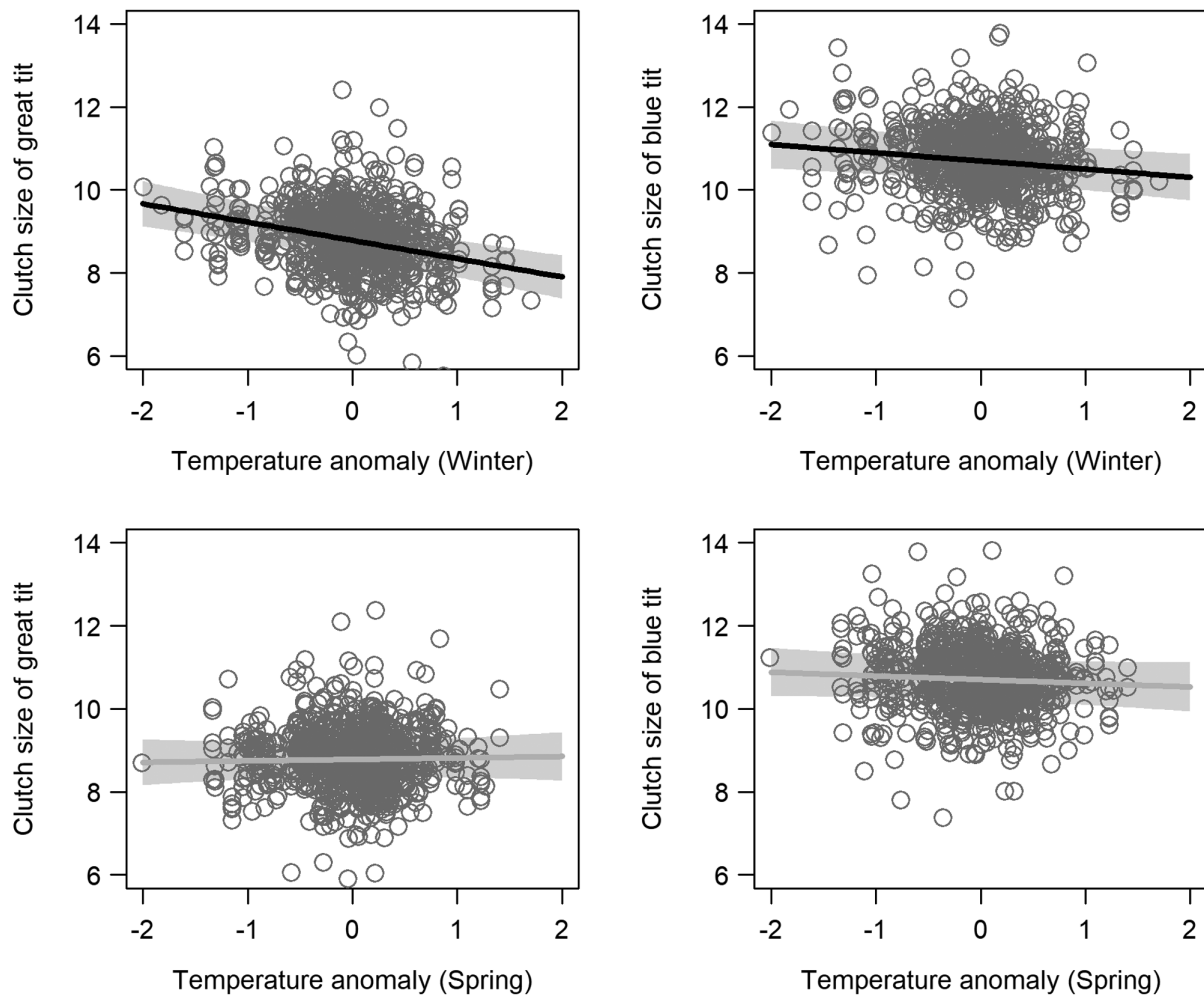


Figure 2. Effects of temperature anomaly ($^{\circ}\text{C}$) in spring and winter on clutch size of great and blue tits. The lines ($\pm 95\%$ CI) are the predicted values obtained from a linear mixed-effect models where latitude, longitude and altitude and nest floor surface are maintained at their mean values and habitat, urbanization and material at their reference values (i.e. conifer, natural habitat and the construction material concrete, respectively). Black lines represent effect sizes that did not include zero in 95% CI and grey lines represent effect sizes that did include zero in 95% CI (see Table 5 and 6 for details).

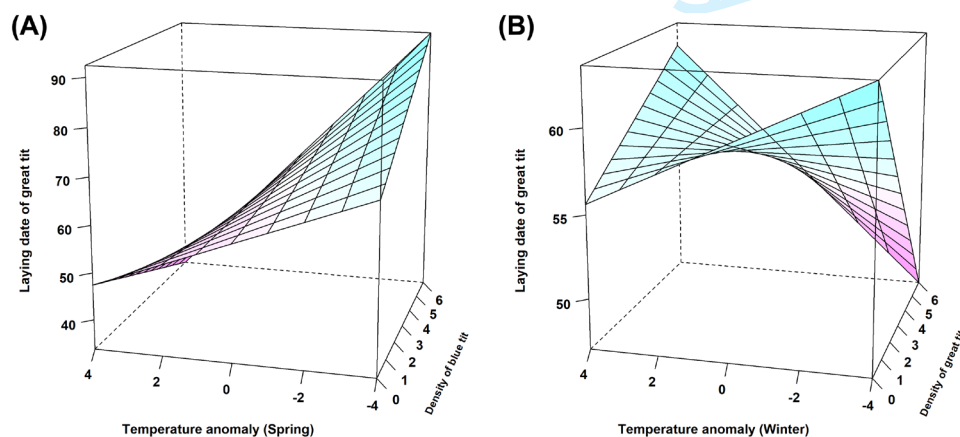


Figure 3. Effects of the interaction term of temperature anomaly ($^{\circ}\text{C}$) in spring and density (no. of nest boxes ha^{-1}) of great tits on laying date of great tits (a) and effects of the interaction term of temperature anomaly in winter and density of great tits on laying date of great tits (b). The surfaces represent the predicted values at average values for other predictors included in the model obtained from a final model after averaging the top models obtained from 32 768 candidate models.

Table 3. Summary results after model averaging of the effects of temperature anomaly in spring and winter and density of great and blue tit on laying date of blue tit across Europe and North Africa. Estimates (SE) and 95% CI are reported. The model also included other potential predictors of laying date of great tit. Sample size is 919 observations on yearly average laying date taken across 75 sites across Europe. Effects in bold font are statistically significant.

Parameter	Estimate	SE	CI (95%)
Intercept	58.836	2.273	(54.374, 63.299)
Density of blue tit (BT)	1.946	0.651	(0.667, 3.224)
Density of great tit (GT)	−1.713	0.519	(−2.732, −0.693)
Latitude	22.628	2.738	(16.475, 28.782)
Longitude	0.079	2.155	(−4.150, 4.309)
Floor surface (nest)	−2.191	1.038	(−4.229, −0.153)
T^a Spring	−4.925	0.526	(−5.959, −3.891)
T ^a winter	−0.469	0.471	(−1.394, 0.454)
T^a Spring × BT	−2.794	0.732	(−4.232, −1.357)
T^a Spring × GT	−1.512	0.654	(−2.796, −0.227)
T ^a Winter × BT	−0.652	0.731	(−2.087, 0.872)
T^a Winter × GT	1.472	0.571	(0.350, 2.594)
Altitude	15.362	2.620	(10.219, 20.506)
Material	1.307	1.907	(−2.436, 5.051)
Urbanization	−2.592	1.080	(−4.713, −0.472)
Habitat (Deciduous)	−5.985	2.626	(−11.139, −0.831)
Habitat (Evergreen)	6.846	3.793	(−0.598, 14.290)
Habitat (Mixed)	−3.684	0.883	(−5.418, −1.951)

* Effect sizes have been standardized to two SD following Gelman (2008). Reference levels were (no) urbanized for urbanization and (conifer) for habitat. Parameter estimates for predictors with confidence intervals not including zero are highlighted in bold. CI is confidence interval.

In contrast, clutch size increased with increasing spring temperature anomaly when blue tit density increased (Fig. 5c). These effects were found while controlling for confounding variables known to affect laying date, such as latitude, longitude, altitude, habitat, urbanization, nest floor surface and nest-box material.

Effects on blue tit clutch size

Opposite to what we found for great tit clutch size, we did not find any interaction effect between temperature anomaly in either spring or winter and the density of either great or blue tit on clutch size of blue tit. These effects were found while controlling for confounding variables known to affect clutch size, such as laying date, latitude, longitude, altitude, habitat, urbanization, nest floor surface and nest box material.

Discussion

An increase in temperature anomaly due to climate warming was correlated with the timing of breeding and clutch size in great and blue tits across large spatial and temporal scales in Europe and North Africa. This is not a novel finding since effects of temperature anomaly on laying date and clutch size have previously been found in these species (Visser et al. 1998, 2003). What is novel though is that the strength of the effects of temperature anomaly on laying date and clutch size varied with density, as shown by multiple significant interaction terms between density and temperature anomaly in both winter and spring. Furthermore, this is the first study showing that the effect of temperature anomaly on laying date and clutch size is widespread across large spatial and temporal scales in these coexisting tits.

In this study we focused on how intra- and interspecific density interacted with climate warming, and how this affected two species of coexisting tits. Blue and great tit advanced laying date as spring temperature increased (prediction 1: supported). In contrast, winter temperature did not correlate with laying date of either great or blue tit (prediction 2: not supported). However, when winter temperature increased both great and blue tits delayed laying date as density increased (prediction 3: supported). Specifically, that was the case for laying date of great tits when intraspecific density increased and for laying date of blue tits when heterospecific density increased. In contrast, when spring temperatures increased both great and blue tits advanced laying date as density increased (prediction 4: not supported). Specifically, it occurs for laying date of great tits when heterospecific density increased, and for laying date of blue tits when both hetero- and conspecific density increased.

We expected that an increase in spring temperature anomalies should advance laying date, and that an increase in density should delay laying, which could cancel the effect of spring temperature anomaly on laying date (prediction 4). As expected, an increase in spring temperature anomaly resulted in a similar advance in laying date in the two tits species. In contrast, sites where density of conspecifics or heterospecifics is high, an additional increase in temperature anomaly in spring resulted in further advance in laying date. It is well known that social information gathered from conspecifics and heterospecifics show that similar ecological niches could serve as important cues to select breeding habitat or to change behaviour related to the acquisition of food, as has been already demonstrated in birds, including great tit (Aplin et al. 2015, Samplonius et al. 2017). For instance, migratory pied flycatcher *Ficedula hypoleuca* could gather social information from the breeding phenology of great tits as a social cue to select its breeding habitat later during their breeding season (Samplonius and Both 2017). It could be possible that the presence of more conspecifics or heterospecifics could act as social cues that serve great and blue tits to advance laying date more to cope with an earlier emergence of caterpillars and hence improving their responses to climate

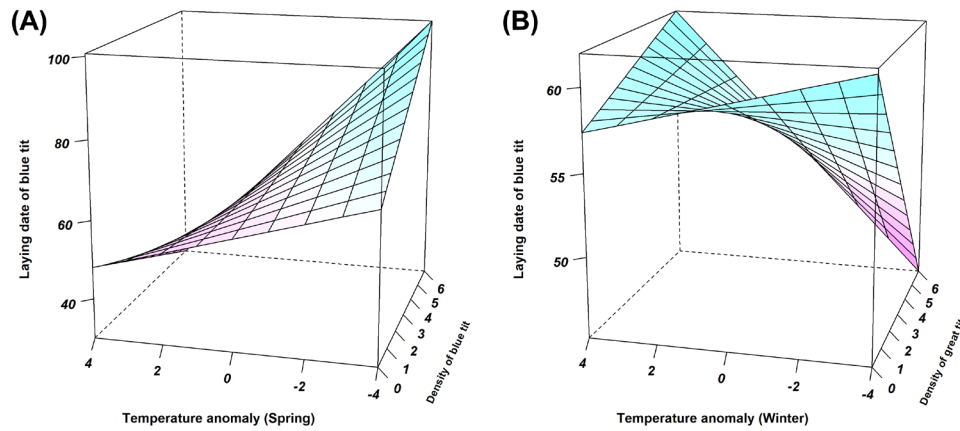


Figure 4. Effects of the interaction term of temperature anomaly ($^{\circ}\text{C}$) in spring and density (number of nest boxes ha^{-1}) of blue tits on laying date of blue tits (a) and effects of the interaction term of temperature anomaly in winter and density of great tits on laying date of blue tits (b). The surfaces represent the predicted values at average values for other predictors included in the model obtained from a final model after averaging the top models obtained from 32 768 candidate models.

Table 4. Summary results after model averaging of the effects of temperature anomaly in spring and winter and density of great and blue tit on clutch size of great tit across Europe and North Africa. Estimates (SE) and 95% CI are reported. The model also included other potential predictors of clutch size of great tit. Sample size is 916 observations on yearly average laying date taken across 75 sites across Europe. Effects in bold font are statistically significant.

Parameter	Estimate	SE	CI (95%)
Intercept	8.642	0.200	(8.247, 9.036)
Laying date of great tit	-0.839	0.094	(-1.024, -0.653)
Density of blue tit (BT)	-0.150	0.102	(-0.351, 0.050)
Density of great tit (GT)	-0.203	0.080	(-0.362, -0.044)
Latitude	0.145	0.404	(-0.648, 0.939)
Longitude	1.110	0.180	(0.759, 1.464)
Floor surface (nest)	0.101	0.143	(-0.179, 0.382)
T ^a Spring	0.040	0.092	(-0.140, 0.222)
T^a Winter	-0.506	0.079	(-0.662, -0.349)
T^a Spring \times BT	0.322	0.122	(0.083, 0.562)
T^a Spring \times GT	-0.277	0.098	(-0.470, -0.084)
T^a Winter \times BT	-0.445	0.120	(-0.682, -0.208)
T ^a Winter \times GT	0.128	0.096	(-0.060, 0.318)
Altitude	-0.480	0.213	(-0.899, -0.060)
Material	0.494	0.208	(0.084, 0.903)
Urbanization	-0.655	0.163	(-0.976, -0.334)
Habitat (Deciduous)	0.339	0.246	(-0.144, 0.824)
Habitat (Evergreen)	-0.211	0.383	(-0.964, 0.541)
Habitat (Mixed)	0.196	0.142	(-0.082, 0.475)

* Effect sizes have been standardized to two SD following Gelman (2008). Reference levels were (no) urbanized for urbanization and (conifer) for habitat. Parameter estimates for predictors with confidence intervals not including zero are highlighted in bold. CI is confidence interval.

change. That would be the case when advanced laying date results in a higher degree of synchrony between emergence of food and timing of breeding.

It is also possible that great and blue tit interact with other migrant or resident species. For instance, it is known that great and blue tits could compete with pied flycatcher *Ficedula hypoleuca*. However, it has been shown that the two species of tit affect pied flycatcher but not the reverse (Wittwer et al. 2015). Although, there could be other interacting species we consider that the most important competitive interaction was recorded in this study. This question could be subject to experimental and observational future studies.

Interestingly, as winter temperatures increased, clutch size declined in both great and blue tits (prediction 6: supported). It should be highlighted, that winter temperature was more strongly correlated with great tit than blue tit since clutch size is reduced more after warmer winters in the former species. In contrast, spring temperatures were not correlated with clutch size in either species (prediction 5: not supported). Climate warming and its interaction with density affected clutch size of great and blue tits differently. Thus, we did not find any significant interaction between winter or spring temperature and density of great or blue tits on clutch size of blue tits. However, that was not the case for great tits since spring and winter temperature interacted with density affecting great tit clutch size. Thus, clutch size of great tit was reduced more when spring temperature and density of conspecifics (i.e. great tits) increased (prediction 7: supported), but it increased more steeply when spring temperature and density of heterospecifics (i.e. blue tits) increased (prediction 7: not supported). Furthermore, winter temperature interacted with competition affecting only clutch size of great tits. Thus, clutch size of great tit was reduced more strongly when winter temperature and density of heterospecific (i.e. blue tits) increased which is in line with results of Dhondt (2010) in which he reports a stronger effect of density on clutch size of great tit than blue tit (prediction 8: supported).

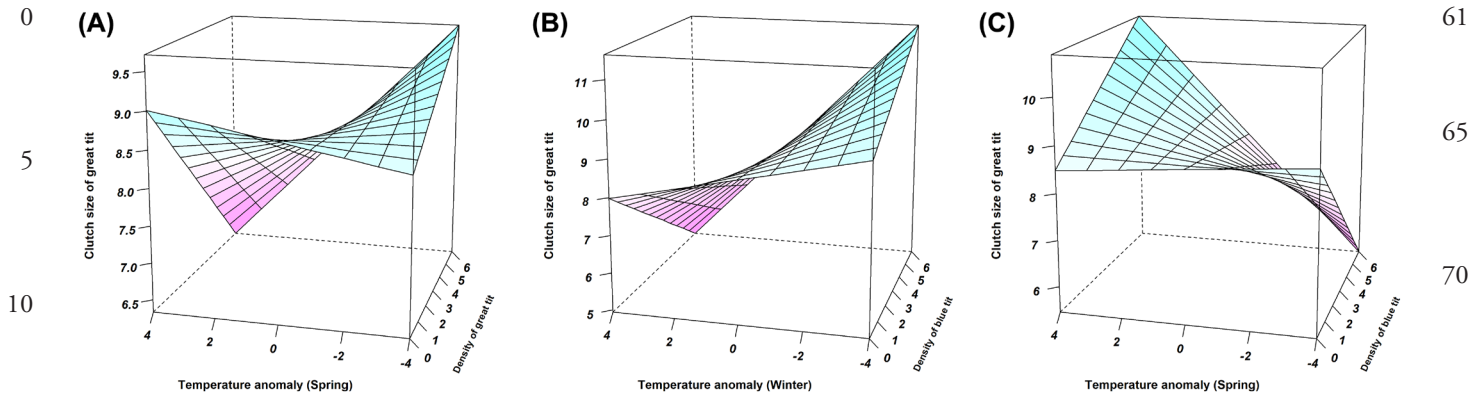


Figure 5. Effects of the interaction term of temperature anomaly (°C) in spring and density of great tits (nest boxes ha⁻¹) on clutch size of great tits (a) and the interaction term of temperature anomaly (°C) in winter and density of blue tits on clutch size of great tits (b) and the interaction term of temperature anomaly (°C) in spring and density of blue tits (nest boxes ha⁻¹) on clutch size of great tits (c). The surfaces represent the predicted values at average values for other predictors included in the model obtained from a final model after averaging the top models obtained from 65 536 candidate models.

Density-dependence could result in a delay in laying date and a reduction in clutch size (Stenseth et al. 2015). An increase in intra- or interspecific competition (resulting from higher densities) could affect the response to climate warming (Stenseth et al. 2015). If that was the case, we would expect a significant interaction between density and temperature anomaly on laying date and clutch size. Indeed, an increase in winter temperature interacted with density of great tit by delaying laying date of both great and blue tit (prediction 3: supported). An increase in winter temperature anomaly could increase adult survival in both species, specifically in blue tits, and this could affect the intensity of competition the subsequent spring, which in turn could reduce clutch size and delay laying at the population level due to density-dependent processes including an increase in the proportion of young inexperienced breeders (Dhondt et al. 1992, Ferrer and Donazar 1996). Importantly, that is what we found for the interactions between winter temperature and density for laying date of great and blue tit and for clutch size of great tit. Alternatively, energetic constraints imposed during winter may also have physiological consequences that may affect laying date. Warm winters may select for metabolic genotypes with a reduced rate of living resulting in smaller clutches (Nilsson and Nilsson 2016). However, our statistical analyses still showed a relationship between temperature anomaly and life history traits after adjustment for density, implying that there are genuine temperature anomaly effects.

Warm springs imply faster caterpillar development, and birds have to speed up to keep up with the caterpillars. One way of doing so is by laying fewer eggs, or by starting incubation earlier (Matthysen et al. 2011). Therefore, it would be possible that the advance in laying date would not suffice to match the emergence of caterpillars, the main food for nestling tits (Perrins 1965, Both et al. 2009). Reduced clutch size may constitute an additional reproductive strategy to decrease the mismatch by advancing incubation, as experimentally shown in the great tit (Bleu et al. 2017). Therefore,

Table 5. Summary results after model averaging of the effects of temperature anomaly in spring and winter and density of great and blue tit on clutch size of blue tit across Europe and North Africa. Estimates (SE) and 95% CI are reported. The model also included other potential predictors of clutch size of great tit. Sample size is 916 observations on yearly average laying date taken across 75 sites across Europe.

Parameter	Estimate	SE	CI (95%)
Intercept	9.705	0.254	(9.205, 10.205)
Laying date of blue tit	-0.954	0.107	(-1.165, -0.743)
Density of blue tit (BT)	-0.143	0.097	(-0.335, 0.047)
Density of great tit (GT)	-0.109	0.080	(-0.267, -0.047)
Latitude	1.560	0.467	(0.643, 2.478)
Longitude	0.491	0.234	(0.030, 0.951)
Floor surface (nest)	-0.282	0.153	(-0.583, 0.017)
T ^a Spring	-0.146	0.079	(-0.301, 0.008)
T^a Winter	-0.159	0.067	(-0.291, -0.026)
T ^a Spring × BT	0.032	0.111	(-0.185, 0.250)
T ^a Spring × GT	-0.136	0.086	(-0.306, 0.034)
T ^a Winter × BT	0.027	0.100	(-0.169, 0.224)
T ^a Winter × GT	0.106	0.093	(-0.077, 0.290)
Altitude	-1.021	0.339	(-1.687, -0.355)
Material	0.608	0.261	(0.096, 1.121)
Urbanization	-0.020	0.168	(-0.350, 0.309)
Habitat (Deciduous)	0.272	0.296	(-0.309, 0.854)
Habitat (Evergreen)	-0.926	0.467	(-1.843, -0.010)
Habitat (Mixed)	0.232	0.142	(-0.047, 0.513)

* Effect sizes have been standardized to two SD following Gelman (2008). Reference levels were (no) urbanized for urbanization and (conifer) for habitat. Parameter estimates for predictors with confidence intervals not including zero are highlighted in bold. CI is confidence interval.

a mismatch between emergence of food and timing of reproduction could also occur at large spatial and temporal scales across the breeding season, as already shown for specific European populations of birds (Visser et al. 1998, Both et al. 2009). In contrast, there is only little evidence of mismatch between blue and great tits and their food resources except for summers with warmer spring (Burgess et al. 2018). Therefore, there are different explanations for the observed reduction in clutch size with increasing temperatures in winter, and this should be explored further in future studies.

Since increasing temperature anomaly in winter reduced clutch size of both species, the abundance of both species should be affected equally by climate warming. However, winter temperature anomaly has a stronger effect on clutch size in great tit (estimate (SE) = -0.506 (0.079), Table 4) than in blue tit (estimate (SE) = -0.159 (0.067), Table 5). However, clutch size is only a component of fitness and we need future studies to try to find out how these contrasting effects translate into differences in population sizes between these tits species. The interacting effect of climate warming with density affected clutch size of great tit while it did not affect clutch size of blue tits. For instance, an increase in spring temperature interacted differently with density of blue tit compared to the effect of density of great tits on their clutch size. Specifically, increasing spring temperature interacted with density of blue tits increasing clutch size of great tit (i.e. rejecting prediction 7), while increasing spring temperatures interacted with density of great tits to decrease clutch size of great tits (i.e. supporting prediction 7). In contrast, the interaction between spring temperature anomaly and density of great or blue tits on clutch size of blue tits was not significant. That is in accordance with prediction 7 since we predicted stronger effects on great compared with blue tits. Therefore, a rapid advance in laying date with density and temperature anomaly did not provide a reproductive advantage for great tits, while that could be the case for blue tits since clutch size was not reduced in blue tits when temperature anomaly and density interacted to affect clutch size. The difference in response to clutch size in great compared to blue tits could be related that either blue tits consume earlier instars of caterpillar larvae that emerge earlier or to that the two species may forage in different micro-habitats, such as different heights in trees with great tits consuming more food on the ground (Slagsvold and Wiebe 2007). The phenology of invertebrate prey may differ between such micro-habitats (Visser et al. 2003, Tremblay et al. 2005), and, therefore, the difference in prey response to climate change by the two species could explain small differences in timing of breeding. Therefore, if these trends in climate warming continue at similar rates, these interspecific differences could translate into differences in abundance or population size favouring blue tit since this species seems to be less affected by climate warming than great tit.

In conclusion, when studying the effects of increasing temperature anomalies on timing of reproduction and clutch size at large spatial and temporal scales, we found that great and

blue tits responded to climate warming by advancing timing of reproduction. However, this advance in breeding date with increasing temperature anomaly did not prevent a reduction in clutch size due to increasing temperature anomalies and increasing densities. In addition, the response to increasing temperature anomaly interacted with density by advancing laying date even more at higher densities. This interaction reduced clutch size in great, but not in blue tits. Further studies are needed to examine how intra- and interspecific relationships interact with temperature anomalies, on timing of breeding and breeding success, which may ultimately affect fitness and population size.

Data availability statement

Data are available from the Dryad Digital Repository: <<http://dx.doi.org/10.5061/dryad.p763611>> (Møller et al. 2020).

Acknowledgements – We would like to warmly thank the hundreds of collaborators and contributors who helped with study plot management, data collection, data management, administration, financial support and scientific discussion. Listing their names individually would most probably provide a biased picture of all of their contributions.

Funding – A. Artemyev acknowledges funding by IB KRC RAS no. 0221-2017-0046 and T. Eeva acknowledges funding by the Academy of Finland (project 265859). This study was funded by research project CGL-2016-79568-C3-3-P (to J. C. Senar), from the Ministry of Economy and Competitiveness, Spanish Research Council.

Author contributions – APM and JB conceived the idea. APM and JB analysed data and APM, JB and AAD wrote the manuscript. APM, AAD, FA, AA, JBa, EB, CB, JcB, JCB, JC, FC, MC, MCh, CC, AD, ME, TE, PNE, JTE, AG, AEG, AGG, LG, IH, IRH, PH, SAH, SJ, AJ, RJ, EK, IK, TL, BL, EL, OL MCM, RM, BM, EM, TDM, SM, CM, MM, RGN, J-ÅN, SGN, ACN, MVN, MO, CSP, RP, IP, VR, HR, HuR, SR, JCS, JTS, LPS, TS, TSo, AS, MJS, JT, PT, AJvN, WW and MML collected data and approved final manuscript.

Conflicts of interest – The authors declare not conflicts of interest.

Permits – The data used in every study plot have the appropriated permits issued for each country or region government.

References

- Aplin, L. M. et al. 2015. Experimentally induced innovations lead to persistent culture via conformity in wild birds. – *Nature* 518: 538–541.
- Alatalo, R. V. and Lundberg, A. 1984. Density dependence in breeding success of the pied flycatcher (*Ficedula hypoleuca*). – *J. Anim. Ecol.* 53: 969–977.
- Balbontín, J. and Ferrer, M. 2008. Density-dependence by habitat heterogeneity: individual quality versus territory quality. – *Oikos* 117: 1111–1114.
- Bartón, K. 2009. MuMIn: multi-model inference. – R package, ver. 0.12.2. <<http://r-forge.r-project.org/projects/mumin/>>.

- Bates, D. and Maechler, M. 2009. lme4: linear mixed-effects models using Eigen and Eigen. – R package, ver. 0.999375-31. <<http://CRAN.R-project.org/package=lme4>>.
- Bleu, J. et al. 2017. Nest box temperature affects clutch size, incubation initiation and nestling health in great tits. – *Behav. Ecol.* 48: 489–501.
- Both, C. 2000. Density dependence of avian clutch size in resident and migrant species: is there a constraint on the predictability of competitor density? – *J. Avian Biol.* 31, 412–417.
- Both, C. et al. 2009. Climate change and unequal phenological changes across four trophic levels: constraints or adaptations. – *J. Anim. Ecol.* 78, 73–83.
- Burgess, M. D. et al. 2018. Tritrophic phenological match-mismatch in space and time. – *Nat. Ecol. Evol.* 2: 970–975.
- Burnham, K. P. and Anderson, D. R. 2002. Model selection and multimodel inference: a practical information-theoretic approach, 2nd edn. – Springer.
- CMB and Crouch, J. 2012. Global surface temperature anomalies: background information. – NOAA NCDC.
- Dhondt, A. A. 2010. Effects of competition on great and blue tit reproduction: intensity and importance in relation to habitat quality. – *J. Anim. Ecol.* 79: 257–265.
- Dhondt, A. A. 2012. Interspecific competition in birds. – Oxford Univ. Press.
- Dhondt, A. A. and Eyckerman, R. 1980. Competition between the great tit and the blue tit outside the breeding season in field experiments. – *Ecology* 61: 1291–1296.
- Dhondt, A. A. and Schilleman, J. 1983. Reproductive success of the great tit in relation to its territorial status. – *Anim. Behav.* 31: 902–912.
- Dhondt, A. A. et al. 1992. Density-dependent clutch size caused by habitat heterogeneity. – *J. Appl. Ecol.* 61: 643–648.
- Ferrer, M. and Donazar, J. A. 1996. Density-dependent fecundity by habitat heterogeneity in an increasing population of Spanish imperial eagles. – *Ecology* 77: 69–77.
- Freund, R. J. et al. 2003. Regression using JMP. – SAS Inst.
- Gelman, A. 2008. Scaling regression inputs by dividing by two standard deviations. – *Statist. Med.* 27: 2865–2873.
- Gibb, J. A. and Betts, M. M. 1963. Food and food supply of nestling tits (*Paridae*) in Breckland pine. – *J. Anim. Ecol.* 32: 489–533.
- Grueber, C. E. et al. 2011. Multimodel inference in ecology and evolution: challenges and solutions. – *J. Evol. Biol.* 24: 699–711.
- Gustafsson, L. and Nilsson, S. G. 1985. Clutch size and breeding success of pied and collared flycatchers *Ficedula* spp. – *Ibis* 127: 380–385.
- Hansen, J. and Lebedeff, S. 1987. Global trends of measured surface air temperature. – *J. Geophys. Res.* 92: 13345–13372.
- Haywood, S. 1993. Sensory and hormonal control of clutch size in birds. – *Q. Rev. Biol.* 68: 33–60.
- Husby, A. et al. 2009. Decline in the frequency and benefits of multiple brooding in great tits as a consequence of a changing environment. – *Proc. R. Soc. B* 276: 1845–1854.
- Kennedy, C. E. J. and Southwood, T. R. E. 1984. The number of species of insects associated with British trees: a re-analysis. – *J. Anim. Ecol.* 53: 455–478.
- Kluijver, H. N. 1951. The population ecology of the great tit *Parus m. major* L. – *Ardea* 39: 1–135.
- Lambrechts, M. M. et al. 1997. A single response mechanism is responsible for evolutionary adaptive variation in a bird's laying date. – *Proc. Natl Acad. Sci. USA* 94: 5153–5155.
- Lambrechts, M. M. et al. 2008. Double-brooded great tits (*Parus major*) in Mediterranean oak habitats: do first broods always perform better than second broods? – *Russ. J. Ecol.* 39: 516.
- Lambrechts, M. M. et al. 2010. The design of artificial nest boxes for the study of secondary hole-nesting birds: a review of methodological inconsistencies and potential biases. – *Acta Ornithol.* 45: 1–26.
- Matthysen, E. M. et al. 2011. Multiple responses to increasing spring temperatures in the breeding cycle of blue and great tits (*Cyanistes caeruleus*, *Parus major*). – *Global Change Biol.* 17: 1–16.
- Møller, A. P. et al. 2010. Effects of climate change on birds. – Oxford Univ. Press.
- Møller, A. P. et al. 2014a. Clutch size in European secondary hole-nesting passerines in relation to nest-box floor area, habitat, geographic location and study year. – *Methods Ecol. Evol.* 5: 353–362.
- Møller, A. P. et al. 2014b. Variation in clutch size in relation to nest size in birds. – *Ecol. Evol.* 4: 3583–3595.
- Møller, A. P. et al. 2018. Effects of interspecific coexistence on laying date and clutch size in two closely related species of hole-nesting birds. – *J. Anim. Ecol.* 87: 1738–1748.
- Møller, A. P. et al. 2019. Effects of climate change on birds. – Oxford Univ. Press.
- Møller, A. P. et al. 2020. Data from: Interaction of climate change with effects of conspecific and heterospecific density on reproduction. – Dryad Digital Repository, <<http://dx.doi.org/10.5061/dryad.p.763611>>.
- Nager, R. G. and van Noordwijk, A. J. 1995. Proximate and ultimate aspects of phenotypic plasticity in timing of great tit breeding in a heterogeneous environment. – *Am. Nat.* 146: 454–474.
- Nilsson, J.-F. and Nilsson, J.-Å. 2016. Fluctuating selection on basal metabolic rate. – *Ecol. Evol.* 6: 1197–1202.
- Parmesan, C. and Yohe, G. 2003. A globally coherent fingerprint of climate change impacts across natural systems. – *Nature* 421: 37–42.
- Perrins, C. M. 1965. Population fluctuations and clutch-size in the great tit, *Parus major*. – *J. Anim. Ecol.* 34: 601–647.
- Perrins, C. M. 1991. Tits and their caterpillar food supply. – *Ibis* 133: 49–54.
- Pearce-Higgins, J. W. and Green, R. E. 2014. Birds and climate change: impacts and conservation solutions. – Cambridge Univ. Press.
- Richards, S. A. 2008. Dealing with overdispersed count data in applied ecology. – *J. Appl. Ecol.* 45: 218–227.
- Robinson, R. A. et al. 2007. Weather-dependent survival: implications of climate change for passerine population processes. – *Ibis* 149: 357–364.
- Samplonius, J. M. and Both, C. 2017. Competitor phenology as a social cue in breeding site selection. – *J. Anim. Ecol.* 86: 615–23.
- Samplonius, J. M. et al. 2017. Nest site preference depends on the relative density of conspecifics and heterospecifics in wild birds. – *Front. Zool.* 14: 56.
- Slagsvold, T. and Wiebe, K. L. 2007. Learning the ecological niche. – *Proc. R. Soc. B* 274: 19–23.
- Smith, R. J. and Moore, F. R. 2004. Arrival timing and seasonal reproductive performance in a long-distance migratory landbird. – *Behav. Ecol. Sociobiol.* 57: 231–239.
- Stenning, M. 2018. The blue tit. – T. and A. D. Poyser, London, UK.

- 0 Stenseth, N. C. et al. 2015. Testing for effects of climate change on competitive relationships and coexistence between two bird species. – Proc. R. Soc. B 282: 20142929.
- 5 Tremblay, I. et al. 2005. The effect of habitat quality on foraging patterns, provisioning rate and nestling growth in Corsican blue tits *Parus caeruleus*. – Ibis 147: 17–24.
- 10 Török, J. and Tóth, L. 1988. Density dependence in reproduction in the collared flycatcher (*Ficedula albicollis*) at high population levels. – J. Anim. Ecol. 57: 251–258.
- 15 Vaugoyeau, M. et al. 2016. Interspecific variation in the relationship between clutch size, laying date and intensity of urbanisation in four species of hole-nesting birds. – Ecol. Evol. 6: 16.
- 20 Welden, C. W. and Slauson, W. L. 1986. The intensity of competition versus its importance – an overlooked distinction and some implications. – Q. Rev. Biol. 61: 23–44.
- 25 Visser, M. E. et al. 1998. Warmer springs lead to mistimed reproduction in great tits (*Parus major*). – Proc. R. Soc. B 265: 1867–1870.
- 30 Visser, M. E. et al. 2003. Variable responses to large scale climate change in European *Parus* populations. – Proc. R. Soc. B 270: 367–372.
- 35 Visser, M. E. et al. 2009. Temperature has a causal effect on avian timing of reproduction. – Proc R. Soc. B 276: 2323–2331.
- 40 Winkler, D. W. and Allen, P. E. 1996. The seasonal decline in tree swallow clutch size: physiological constraint or strategic adjustment? – Ecology 77: 922–932.
- 45 Wittwer, T. et al. 2015. Long-term population dynamics of a migrant bird suggests interaction of climate change and competition with resident species. – Oikos 124: 1151–1159.

Author Queries

JOB NUMBER: 7305

JOURNAL: OIK_OIK

- Q1 Figures must be cited in strict sequential order, in the text the figures are cited out of order: please choose whether to re-number the figures or remove this citation.

For Review Only